

## A phase III, single-arm study to evaluate the efficacy and safety of ONCOFID-P-B (paclitaxel-hyaluronic acid conjugate) administered intravesically to patients with BCG- unresponsive Carcinoma in Situ of the bladder with or without Ta-T1 papillary disease

**Status:** Recruiting

### Eligibility Criteria

**Sex:** Male or Female

**Age Group:** 18 years and over

This study is NOT accepting healthy volunteers

**Inclusion Criteria:**

- persistent or recurrent confirmed carcinoma in situ (CIS) of the bladder - unresponsive to BCG treatment and refuse radical cystectomy or are not clinically suitable for cystectomy - able to walk and capable of all selfcare but unable to carry out any work activities; up and about more than 50% of waking hours - women and men of child bearing age must follow specific requirements for birth control

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**Exclusion Criteria:**

- current or previous muscle-invasive cancer or metastatic urothelial cancer - current or prior systemic therapy for bladder cancer. - women who are pregnant or breast feeding - additional medical or mental health diagnosis (study staff will review)

### Conditions & Interventions

**Interventions:**

Drug: ONCOFID P-B (PACLITAXEL-HYALURONIC ACID)

**Conditions:**

Cancer

**Keywords:**

Clinics and Surgery Center (CSC), Bladder Cancer in Situ (CIS), Bladder CIS

### More Information

**Description:** The purpose of this study is to understand if the study medication ONCOFID-P-B is effective and safe in treating patients with carcinoma in situ of the bladder who have not received benefit from standard BCG treatment and are not candidates for radical cystectomy.

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